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NEWS NEWS	1 2	NOV	21	Web Page for STN Seminar Schedule - N. America CAS patent coverage to include exemplified prophetic substances identified in English-, French-, German-,
	_			and Japanese-language basic patents from 2004-present
NEWS		NOA		MARPAT enhanced with FSORT command
NEWS	4	NOV		CHEMSAFE now available on STN Easy
NEWS	5	NOV	26	Two new SET commands increase convenience of STN searching
NEWS	6	DEC	01	ChemPort single article sales feature unavailable
NEWS	7	DEC		GBFULL now offers single source for full-text coverage of complete UK patent families
	_	DEC		
NEWS NEWS	8	JAN		Fifty-one pharmaceutical ingredients added to PS
	-			The retention policy for unread STNmail messages will change in 2009 for STN-Columbus and STN-Tokyo
NEWS	10	JAN	07	WPIDS, WPINDEX, and WPIX enhanced Japanese Patent Classification Data
NEWS	11	FEB	02	Simultaneous left and right truncation (SLART) added for CERAB, COMPUAB, ELCOM, and SOLIDSTATE
NEWS	12	FEB	0.2	GENBANK enhanced with SET PLURALS and SET SPELLING
NEWS		FEB		Patent sequence location (PSL) data added to USGENE
NEWS				COMPENDEX reloaded and enhanced
NEWS				WTEXTILES reloaded and enhanced
NEWS		FEB		New patent-examiner citations in 300,000 CA/CAplus
Hano	10		1,	patent records provide insights into related prior art
NEWS	17	FEB	19	Increase the precision of your patent queries use terms from the IPC Thesaurus, Version 2009.01
NEWS	18	FEB	23	Several formats for image display and print options discontinued in USPATFULL and USPAT2
NEWS	19	FEB	23	MEDLINE now offers more precise author group fields
NEWS	20	FEB	23	and 2009 MeSH terms TOXCENTER updates mirror those of MEDLINE - more
				precise author group fields and 2009 MeSH terms
NEWS	21	FEB	23	Three million new patent records blast AEROSPACE into STN patent clusters
NEWS	22	FEB	25	USGENE enhanced with patent family and legal status
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chain nodes :
6 7 8
ring nodes :
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ring/chain nodes :
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chain bonds :
1-6 2-7 3-8 5-15 15-16
ring bonds :
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exact/norm bonds :
1-2 1-5 1-6 2-3 2-7 3-4 3-8 4-5 5-15
exact bonds :
10-11 10-14 11-12 12-13 13-14 15-16
isolated ring systems :
containing 10 :
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G1:C,O,S,N

Match level: 1:Atom 2:Atom 3:Atom 4:Atom 5:Atom 6:CLASS 7:CLASS 8:CLASS 10:Atom 11:Atom 12:Atom 13:Atom 14:Atom 15:CLASS 16:CLASS 19:CLASS

50 ANSWERS

L1 STRUCTURE UPLOADED

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L1 HAS NO ANSWERS

L1 STR



G1 C, O, S, N

Structure attributes must be viewed using STN Express query preparation.

6912 TO

9328

=> S L1

SAMPLE SEARCH INITIATED 08:19:18 FILE 'REGISTRY'

SAMPLE SCREEN SEARCH COMPLETED - 1384 TO ITERATE

100.0% PROCESSED 1384 ITERATIONS

INCOMPLETE SEARCH (SYSTEM LIMIT EXCEEDED) SEARCH TIME: 00.00.01

FULL FILE PROJECTIONS: ONLINE **COMPLETE**

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PROJECTED ITERATIONS: 25449 TO 29911

L2 50 SEA SSS SAM L1

PROJECTED ANSWERS:

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=> S L1 FULL

FULL SEARCH INITIATED 08:19:33 FILE 'REGISTRY'

FULL SCREEN SEARCH COMPLETED - 27013 TO ITERATE

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L3 7216 SEA SSS FUL L1

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=> S L3 1.4

192 T.3

=> D IRIR ARS HITSTR 180-192

14 AREMER 180 OF 192 CAPLUS COPYRIGHT 2009 ACS on STN ACCUSSION NUMBER: 1960:44804 CAPLUS DOCUMENT NUMBER: 54:44804 CAIGURAL REFERENCE NO.: 54:8791F-h

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of appropriate aldebyds in the presence of StOMs is AcOS 2 hrs. gave the following deriver: 1-thinamylidese, DVI n. 229-277 [8].

201-14. Takes treated with dry NUI in Mexico columns gave NUE thoisinks—1-setter, 371, decomposed 391-27, 5-bentylidese derivative, 85%, decomposed 399-77; 5-in-Aritownzylidese derivative, 85%, decomposed 399-77; 5-in-Aritownzylidese) derivative, 95%,

184, decomposed new Fr Primanassempassement (1988). The Composed 199-67 (1984). The Composed 199-67 (1984) in Composed 200-97 (1984) in Composed

NN 112715-36-3 CAPLUS CN 3-Thisrolidinescetic Thiarolidineacetic acid, 5-(2-furanylmethylene)-4-oxo-2-thioxo-, monium malt [1:1] (CA INDEX NAME)

L4 ANSMER 180 OF 192 CAPLUS COPYRIGHT 2009 ACS on STN (Continued) IT 99073-34-4, 3-Tharolidaseactic acid,
5-fm fmyliders-t-oux-2-thicoxN 99073-44 CMRUNS
This olidiseacetic acid, 5-[2-fmranylmethyless-4-oux-2-thicoxINDEX NUMBER.

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L4 AREMER 182 OF 182 CAPLUS COPYRIGHT 2008 ACS on STN ACCESSION NUMBER: 1840:2178 CAPLUS DOUBLENT NUMBER: 54:2178

DOCUMENT NUMBER: 5-11.19 ORIGINAL REFERENCE NO.: 54:490e-q

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14 ANSWER 187 OF 192 CAPLUS COPYRIGHT 2009 ACS on STN (Continued)
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L4 AREMER 183 OF 192 CAPLUS COPTRIGHT 2009 ACS on STN ACCESSION NUMBER: 1950:50558 CAPLUS DOCUMENT NUMBER: 52:50558 ORIGINAL REFERENCE NO.: 52:90626-F Synthesis of derivatives of thiazolidone having biological interest. VII- Synthesis of N-substituted derivatives of rhodanine starting with deconvertate

solution of the control of the contro INGAMORICS: AUTROR(S): CORPORATE SOURCE: SOURCE;

Kb: PKSF (respectation of) (preparation of) 99972-49-3 CAPUJS 4-Thiasolidinome, 5-(2-furanylmethylene)-3-(2-propen-1-yl)-2-thioxo- (CA

14 Demand Ref of 12 Octave Contrainer 1609 Act on 1780
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cooling,
a yellow oil separated The aqueous layer was washed with CRC13 and the extract combined with the oil. After drying with anhydrous Na2504 and

evaporation, 103 g, of a heavy amber oil, 3-(2-hydroxyethyl)rhodanine (I) was obtained. By replacing ethanolanine with equivalent weight of propanolanine or solamine, 3-(3-hydroxypropyl)-and 3-(4-hydroxybutyl)rhodanine, resp., were

Inco.

Equinol. ants. of I and various aromatic and heterocyclic aldehydes were refluxed for 0.5-5 hr. with piperadine or EtN as catalyst to gave 3-(2-)ydroxythyl)-5-(4-diserthylamicobenzyldidens) khodanane, zed-vaolet,

317, 3:[-]-pricopythyl)-5-[4-esthosybensylidene] hhodanime, yellow, 137, 3:[-]-pricopythyl)-5-[4-esthosybensylidene] hhodanime, yellow, 132-137, 3:[-]-pricopythyl]-3-[-]-pricopylidene] hhodanime, ocange, m. 132-137, 3:[-]-pricopythyl]-3-[-]-intaintobensylidene] hhodanime, ocandomenylidene] hodanime, yellow, m. 132-607, 3:[-]-pricopylidene] hodanime, yellow, m.

224-57, 2-(2-bydrovyethyl)-5-furfurylidenerhodanne, yellow, n. 150-57, 3-(2-bydrovyethyl)-5-(2-ethorybensylidene) thodanne, 150-57, 3-(2-bydrovyethyl)-5-(2-ethorybensylidene) thodanne, 150-57, 3-(2-bydrovyethyl)-5-(2-ethorybensylidene) pyriddylidenylodanna, yellow, n. 177-57. These compdises were condensed with 1st styrene-malcic anhytride oppolyment by heating in pyridise nolution for 2-3-5 has to days inject-annitive remain usefur the pyridise nolution for 2-3-5 has to days inject-annitive remain usefur

lathographic plates.
99185-08-7, Nbdanne, 5-furfurylidese-3-(2-hydronyethyl)Ireartion with maleic ambydride copolymers, and light-sensitive resins

therefroe)
59183-08-7 CMPUS
6-Thiarolidinose, 5-(2-furarylmethylese)-3-(2-hydroxyethyl)-2-thioxo-

10/534,919 03/05/2009 14 ANSMER 185 OF 192 CAPLUS COPYRIGHT 2009 ACS on STN ACCESSION NUMBER: 1958:35201 CAPLUS

14 AMENUS 184 OF 192 CAPLUS COPYRIGHT 2009 ACS on STN

ANGUAGE: Unavailable
AB cf. C.A. 50, 14713d. To 67.0 g. SC(SCE2CO2B)2 (I) in 1.2 1. E20 war selected 100 11.5 p. NaCOSO 10 700 AL RED Colleges over 3 her sp 14.4 g. 3 mth/1-demokerachinoste in 500 AL IN 100 ATT 12.15 hr. of the 10.21 hr. of the 10.21

on a steam bath 2.85 g. CS2, the mixture heated 15 min. longer, treated

on a reason bath 2.0% of CHE, the minimum beated 12 min. lower, tracted 2 min. CHE 1000 cross colors are considered as a color of the colors and colors are colors and colors and colors are colors and colors and colors are colors and colors are colors and colors and colors and colors and colors are colors and colors and colors and colors and colors and colors are colors and colors and colors and colors and colors are colors and colors an

after refluxing 2-3 hrs. a precipitate of 85-98% appropriate

after refluxing 2-3 hr. a presipatate of fit-94 appropriate gravita. The president fit of the

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L4 ANSMER 186 OF 192 CAPLUS COPYRIGHT 2X ACCESSION NUMBER; 1956:12278 CAPLUS DOURNEAT NUMBER; 50:12278 GRIGIMAL REFERENCE NO.: 50:2547d-g TITLE: This coldingones. I.

59:2547d-q Thiatolidinones. I. 2-[1-8-8phthylimino]-4-thiatolidinone and its 2-(1-8-8phthylimino]-4. V. Ravenshaw Coll., Cuttack Journal of Scientisio & Industrial Research (1955), 168, 16-18 COEDEM, 93:194C/ 18881 0022-4456 AUTEOR(S): CORPORATE SOURCE: SOURCE:

DOCUMENT TITES

ORDER 037306; 1898 0022-4456

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AND 154* (Proposed by Technological Control Contr

owt in glacial ACOS in the presence of anystross Smooth and also in alc. (95%)
MOST. The mitross derives were prepared in AcOO. The following are the compute. condensed with I (carbonyl compound and m.p. of 5-arylidene compound.

p-decipid-captions-2-1-capthylumino-i-chikanicimous were prepared
treatment of the Scaptiones-2-1-table INE, following West Liebbyde
and Control of the Scaptiones-2-table INE, following West Liebbyde
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Vormal

Vormal I,
m. 191* (from ECOH). A typical condensation, a methylation, and a
reduction are described: 3 g. I was refluxed 1.5 hrs. with 1.5 g. NCHO 30 oc. glacial AcOH and 4 q. dry AcONa, and the mixture poured into N20 yield p-MeCSH4N:C.S.C(:CHR).CO.NH (II); 1 q. II added to 1 q. NCH in 8 EtOH was treated slowly with 3 g. NeI and heated 3 hrs. on a H2O bath until neutral to litnus, to yield on cooling p-MecGHANC.S.C1(CHS).CO.NMc [III); finally, treating 1 g. II suspended in 8 ce. H2O containing 1.5

14 AMENUR 188 OF 192 CAPLUS COPYRIGHT 2009 ACS ON STN

L4 ANSMER 189 OF 192 CAPLUS COPTRIGHT 2009 ACS on STN ACCESSION NUMBER: 1955:79479 CAPLUS DOZUMENT NUMBER: 49:79479

ORIGINAL REFERENCE NO.: 49:7556b-q

49:7550-q
3-71:nbhorosethanesulfenyloxarolidine- and
thiazolidine-2,4-diomes
Crosall, M. 3,7 Lo, Chien-Pen; Shropshire, Elwood Y.
Roha & Basa, Philadelphia, IM.
Soursal of the American Chemical Society (1953), 75,
CODE, UNCERT, ISSN: 0002-7863
WORTHSH (1958), 10002-7863

DOCUMENT TYPE:

DOUBMENT TITLE

OF THE PROPERTY OF THE PROPERT

xi A: (Bu glycolate (72 g.), 32 g. urea, 29 g. NaONe, and 250 cc. absolute

Etcs $_{\rm Elowly}$ serated and refluxed 2 h., the Etcs distilled in vacuo, 200 cc. water added and removed in varue, the cooled colution of the 3a salt of coarchidms-2,4-drome treated with 100 g. CliCCL in 100 cc. part, other, other, 3-dribble consistency of the column of the coarchidms-2,4-drome, an 18-20 g. 3-dribble consistency of the coarchidms-2,4-drome, an 18-20 g. Section 100 cc. water coarching 2,4-drome, and 1-drome coarching 2,4-drome coarching 2,4-drome, and 1-drome coarching 2,4-drome coarching

mixture stirred several hrs. at room temperature, the aqueous layer sated with COL4, and the components of the wave yielded 37 g. and the combined COL4 poline, evaporated in wave yielded 37 g. and the combined COL4 poline in the cold poline in the cold

method, & yield, m.p. (uncor.) given): No. H. A. 70, oil; No. No. A.

Section 9, 19 and 19, 1000cc, 9 areas over a no no not seen, no no not seen, no 19, 297 kg, New, No, NS, Section 7, 100 following Syd-dissortiumed trichilocomethous multiways:2-4-chiaorolidisediones were also prepared in No. No. 19, No. 1

COMMENT TIPE

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aliphatic ketones containing a thiol ether group. The diomothiarolidine derivs. care detained at the care group. The association are actions obtained at the characteristic and the characteristic acceptance, but an expectation that the cross of activity within the two arries is the same. The most effective computs, were 5-1p-oblorobennylisenes/hodealmes and 5-thenylisenes/basica. Which is 18 concentration on cloth year 154 or

C-templiformibonizate, which is 1 memorization on cloth year 15 or it sensitive the control of t

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COCCOMM Sancher, A. Gomer, Permander-Bolanos, J. Univ. Seville Anales de la Beal Sociedad Espanola de Fisio Quinica, Serie B: Quinica (1953), 49B, 51-6 ODDR: ASONA, 1850: 0034-080X CORPORATE SOURCE: DOUBLET TIPS:

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SINCE FILE TOTAL ENTRY SESSION 80.82 267.88

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chain nodes :
6 7 8 23
ring nodes :
1 2 3 4 5 10 11 12 13 14 17 18 19 20 21 22
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chain bonds :
1-6 2-7 3-8 5-15 11-15 14-17
ring bonds :
1-2 1-5 2-3 3-4 4-5 10-11 10-14 11-12 12-13 13-14 17-18 17-22 18-19
19-20 20-21 21-22
exact/norm bonds :
1-2 1-5 1-6 2-3 2-7 3-4 3-8 4-5
exact bonds :
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normalized bonds :
17-18 17-22 18-19 19-20 20-21 21-22
isolated ring systems :
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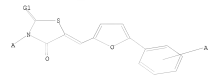
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Match level :

1:Atom 2:Atom 3:Atom 4:Atom 5:Atom 6:CLASS 7:CLASS 8:CLASS 10:Atom 11:Atom 12:Atom 13:Atom 15:CLASS 17:CLASS 18:Atom 19:Atom 20:Atom 21:Atom 22:Atom 23:CLASS 24:Atom

L5 STRUCTURE UPLOADED

=> D L5 HAS NO ANSWERS L5 STR



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Structure attributes must be viewed using STN Express query preparation.

=> S L5 FULL SUB=L3
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FILE 'CAPLUS' ENTERED AT 08:29:36 ON 05 MAR 2009
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FILE COVERS 1907 - 5 Mar 2009 VOL 150 ISS 10 FILE LAST UPDATED: 4 Mar 2009 (20090304/ED)

Caplus now includes complete International Patent Classification (IPC) reclassification data for the third quarter of 2008.

CAS Information Use Policies apply and are available at:

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This file contains CAS Registry Numbers for easy and accurate substance identification.

=> S L6 L7 76 L6 => D IBIB ABS HITSTR 70-76 LT ARSMER TO OF TO CAPLUS COPYRIGHT 2009 ACS on STN ACCESSION NIMBER: 2000:394190 CAPLUS DOCUMENT NUMBER: 133:30722

heterocyclylmethylene thiazolidizediones and analogs as tumor necrosis thiazolidizediones and analogs as tumor necrosi factor inhibitors Wang, Jing; Earnarayan, Ealyanaranan; Rideout, Mone, Seymour; Thu, Hengyi; Niemeyer, Christina; Ready, Thomas P. Structural Biogniformatics Inc., USA NCT Int. Appl., 127 pp. COURSE, FIEREZ Patent

PATENT ASSISSEE(S): SOURCE:

DOCUMENT TIPE: LANGUAGE: FAMILY ACC. NEW, COUNT: FATENT INFORMATION:

PATERT NO. KIND DATE Al 20000608 NO 1999-US28856 US 1998-206108 A 19981204

US 1999-316415 A 19990521

OTHER SOURCE(S): MARPAT 133:30722

AS The title compds. (I) [wherein NI-W5 together = aliphatic, heterocyclic,

of before a control of the control o

LT ANSWER TO OF TO CAPLUS COPYRIGHT 2009 ACS on STN (Continued)

306732-62-7 CAPLUS 4-Thiarolidinome, 3-(3-methoxypropyl)-5-[[5-(2-mitrophenyl)-2-furanyl]methylamej-2-thioxo- (CA INDEX NAME)

N13663-20-6 CAPLUS |-Thiazolidinome, J-(2-propen-1-y1)-2-thioxo-5-[[5-[J-|trifluoromethy])pheny1]-2-furany1]nethylene|- (CA INDEX NAME)

NN 313663-45-5 CAPLUS CN 4-Thiarolidinome, 5-[(5-[2-chlorophenyl)-2-furanyl]methylene]-3-[2-propen-1-yl]-2-thiaxo- (CA INDEX NAME)

324070-85-1 CAPLUS 4-Thiarelaidinese, 5-[|5-(2-ehlorophenyl)-2-furanyl]methylene]-3-|(tetrahydro-2-furanyl)methyl]-2-thioxo- (CA INDEX NAME)

L7 ANSMER 70 OF 76 CAPLUS COPYRIGHT 2009 ACS on STN (Continued) superfamily for the prophylaxis and treatment of inflammatory director

data), 247067-90-9 292076-05-2 299904-95-3 306732-62-7 313663-20-6 313663-45-5 324070-85-1 324564-56-5 30395-73-4 331736-73-3 333393-14-9 909790-92-7

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100537-7-1 100537-9-4 110053-9-4 110054-09-9

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HN 292076-05-2 CAPLUS CN 4-Thiarolidinese. 5-[[5-[2,4-dichlorephenyl]-2-furanyl]nethylene]-3-ethyl-2-thioxo- (CA INDEX NAME)

299904-95-3 CAPLUS Benzolo soid, 3-[5-[[4-oxo-3-[2-propen-1-y1)-2-thioxo-5-thiazolidinylidene]nethyl]-2-furanyl]- (CA INDEX NAME)

1.7 ANSWER TO OF THE CAPLUS COPYRIGHT 2009 ACS on STN

RN 324564-45-6 CAPLUS CN 4-Thiarolidinome, 5-[(5-12,5-dish)orophenyl)-2-furanyl]methylene]-3-ethyl-2-thion- (CA INDEX NAME)

330985-73-4 CAPLUS 4-Thiarolidinose, 5-[[5-(5-chloro-2-methylphenyl)-2-furanyl]methylene]-3-(2-propen-1-yl)-2-thioso- (CA INDEX NAME)

331736-73-3 CAPLUS 4-Thiarolidinose, 5-[5-(2-chloropheny1)-2-furany1]methylene]-3-methyl-2-

PAGE 1-A

IN 1100594-21-5 CAPLUS
CN Cycloberanecarboxamide, N-[5-][5-(3-mitrophenyl) ono-2-thioxo-3-thiazolidinyl)- (CA INDEX NUMB)

PAGE 2-A

LT ARSMER TO OF TO CAPLUS COPYRIGHT 2009 ACS on STN 17 AMSMER 70 OF 76 CAPLUS COPTRIGHT 2009 ACS ON STN (Continued)

PAGE 1-A

PAGE 2-A

110094-22-6 CAPIJS Cyclobexane-carboxanide, N-[4-oxo-5-[[5-[4-[[phenylnethyl)thio]phenyl]-2-furanyl]methylene)-2-thioxo-3-thiarolidinyl]- (CA INDEX NAME)

PAGE 1-A

17 1775-4-02 17775-4-02 17775-9-0-0
17775-1-02 17775-1-

277130-38-6 CARLON 3-Thiascoldanehovanoio soid, [[5-(2-natrophenyl)-2-Duranyl]nethylene]-4-oxo-2-thioso-, l-methylethyl ester, (5E)- (CA INDEX NAME)

273730-98-7 CAPLDS 4-Thiarolidisces. 5-[[5-(2-chlorophenyl)-2-furaryl]methyleme]-3-ethyl-2-thicaco-, [583: (CA. 1885K 1988S) Double bond geometry as shown

ANSWER TO OF THE CAPLUS COPYRIGHT 2009 ACS on STN

273731-04-7 CAPLUS 4-Thiarolidinose, 3-(2-furanylmethyl)-5-[[5-(2-nitrophenyl)-2-furanyllnethylane]-2-thioxo-, (5%)- (CA IMBE SUAM) bond geometry as shown.

273733-07-0 CAPLUS 4-Thiazolidinome, 3-ethyl-5-[[5-(3-nitrophenyl)-2-furanyl]methylene]-2-thioxo-, (5E)- (CA INDEX NUME)

273731-25-2 CMPLOS 4-Thiarolidinome, 3-ethyl-5-[[5-(2-nitrophenyl)-2-furanyl]nethylene]-2-thiomo- (CA INDEX NAME)

2 271712-33-2 CAPLUS 2 3-Thiarolidinebecanoto acid, :[[5-[2-mitropheny1)-2-furany1]methyleme]-4-oxo-2-thioxov, 1-methylethyl ester (CA INDEX NAME)

ASSESSED TO OF THE CAPLUS COPYRIGHT 2009 ACR on STN

27373]-60-5 CAPLUS 4-Thiarolidinone, 3-ethyl-2-thioxo-5-[[5-[2-(triflworomethoxy)phenyl]-2-furanyl)nethyleze)- (CA INDEX NME)

REFERENCE COUNTY THERE ARE 1 CITED REPERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE ANSMER TO OF TO CAPLUS COFFEIGHT 2009 MCS on STN (Continued) 2777731-44-5 CAPLUS 4-Thiarolidinose, 3-(2-Curanylmethyl)-5-[[5-(2-nitrophenyl)-2-turanylmethylene]-2-thiosoc (CA INDEX MOME)

273731-47-8 CAPLOS 4-Thiazolidinose, 3-ethyl-5-[[5-(3-nitrophenyl)-2-furanyl]nethylene]-2-thicoso- (CA INDEX NAME)

273731-52-5 CAPLUS 4-Thiasolidimome, 3-methyl-5-[[5-(3-mitrophenyl)-2-furanyl]methylene]-2-thicoco-(CA INDEX NAME)

273731-53-6 CAPLUS 4-Thiazolidinose, 3-methyl-5-[[5-(2-mitrophenyl)-2-furanyl]methylene]-2-thiazon- (CA THREN NAME)

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DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

| No. JP 2000-565894 UB 2001-976949 UB 2003-366796 UB 2004-829864 US 1998-97476P IE, SI, L7
JP 200252371
U8 2002052396
U8 20030195213
U8 20040198741
PRIORITY APPLE. INFO.: US 1998-113212P p 19981222 NO 1000-1107000

Flaviviridae family by administering certain rhodanise derivs., and

- JANUAL 1 OF M. CALUE COTTAINT 2009 ACS ON SEM. (Continued) statiogs thereof, the and tetrapytic relocation attacks acids and shodaine bestoic entid being particularly effective. 210811-63-69 230912-35-89 230912-56-19 AL NO. (Biological activity or effector, except adverse); 800
- IL: NU: (Niclogical activity or effector, except adverse); NSU (Nicological) (Nicological) (Nicological) (Nicological); NSU (Nyahotic preparation); TSU (Threspectic use); NICL (Nicological study); PNUD (Preparation); USAS (Dear) (Phodatine dariver, preparation), compact, and methods for treating or preparation); NICOLOGICAL (NICOLOGICAL ACTIVATION); NICOLOGICAL (NICOLOGIC
- area; 259811-62-6 CAPLUS 3-Thiarolidimeacetic ecid, 5-[[5-[2-chloro-5-(triflworomethyl)phanyl]-2-furanyl|nethylomo|-6-oxo-2-thioxo- (CA INDEX NAME)

259812-53-8 CAPLUS 3-Thiazolidinepropanoic acud, 5-[[5-(3,4-dichlorophenyl)-2-furanyl]nethylene]-4-oxo-2-thicac- (CA INDEX NAME)

259812-54-9 CAPLUS 3-Thiazolidimehexanous acid, 5-[[5-(3,4-dichlorophenyl)-2-furanylinethylene]-4-oxo-2-thioxo- (CA INDEX NAME)

- ANDMER TI OF TE CAPRUS COPPRIGHT 2009 ACS on STN (Continued)
 226T4-28-6 CAPRUS acid,
 3-Thiatolikaproparoic acid,
 ([5-44bcceopheayl)-2-fuszy]imethylene]-4oun-4-thiory (CAINERS MICHAEL)

259811-52-4 CAPLUS
3-Thiarolidinepropanoic acid,
5-[2-chloro-5-(traffluoromethyl)phenyl]-2furanyl]methylene]-4-oxo-2-thioxo- (CA INDEX NAME)

2 259813-61-5 CAPLUS 2 3-Thiarolidimeacetic acid, [[5-13,4-dichlorophenyl)-2-furanyl]methylene)-4-oxo-2-thioxo- (CA INDEX NAME)

25981]-63-7 CAPLUS 3-Thiaroludusecetic acid, 4-oxo-2-thioxo-5-[[5-[3-[txillworesthyliphemyl]-2-furanyl]methyleme]- (CA INDEX NAME)

ANSMER 71 OF 76 CAPLUS COPTRIGHT 2009 ACS on STN (Continued)

212774-8-4 (25921-15-4 25921-6-5 19821-6-7 (29211-6-4 25921-6-5 25981-6-7 (25981-6-7 25981-7-6-25981-7-6 (25981-7-7 25981-7-6-7 1981-7-7 (25981-7-7 25981-8-7 25981-8-7 25981-7-7 25981-7 25981-7 25981-7 25981-7 25981

ogical study, unclassified); TBU (Therapeutic use); BIOL (Exological study);

ANSWER 71 OF 76 CAPLUS COPYRIGHT 1909 ACS on STN (Continued)

259811-65-9 CRMLUS 3-Thiarolidineacette acid, 5-[[5-(4-chlorophenyl)-2-furanyl]nethylene]-4-oxo-2-thioxo- (CA INDEX NAME)

9811-67-1 CAPLUS Thiasolidimeacetic acid, 5-[[5-[2-chloro-5-(trifluorosethyl)phenyl]-2-ramyl]nethjene]-a-methyl-4-ouc-2-thiouc- (CA INDEX NAME)

NN 259811-69-3 CAPLUS

ARREST TI OF 76 CAPLUS COPYRIGHT 2009 ACS on STN (Continued) 3-Thirsolidineproparoic acid, 5-[15-[3,5-bis/trifluoroesthyl)phemyl]-2-teranylimethylenel-4-coro-2-thicor- (CA INDEX NAME)

59811-74-0 CAPLUS "Thiarolidimepropanoic acid, 5-[[5-(3,5-dichlorophenyl)-2-uranylmethylemel-4-coco-2-thioxo- (CA INDEX NAME)

259811-75-1 CAPLUS 3-Thiazolidinepropanoio acid, 5-[[5-(3,5-dimethylphenyl)-2-furanyl]nethylene]-4-coro-2-thicoro- (CA INDEX NAME)

259812-16-3 CAPLOS Beniodo acid, 6-(15-(15-(2-chloro-5-(trifluoromethyl)phenyl)-2-furamyl)methylene)-6-oxo-2-thioxo-3-thiazolidinyl]methyl)- (CA INDEX

LT AMMER TI OF TO CAPLUS COPYRIGHT 2009 ACS on STN

REPERENCE COUNTY THERE ARE 6 CITED REPERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE TOTAL

DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

PATERT NO. KIND DATE APPLICATION NO. JP 11302280 PRIORITY APPLN, INFO.: OTHER SOURCE(S): MARPAT 131:299442

The title compds: I [A1, A2, A7 = 0, 8; R1 = alkyl, alkenyl, etc.; R2 = alkyl, etc.; m=1-7; Eing B1= beteropyelic Eing, etc.; dotted line indicates zingle or double bond] are prepared. In an in vitro test using BI=0 cells, the title compound II at 3 μ M gave 100% inhibition of

AREMER 72 OF 76 CAPLUS COPYRIGHT 2009 ACS on STN (Continued) RL: BAC (Biological activity or effector, except adverse); ESU

ological study, unclassified; DN (Gymthetic proparation); DN (Therapouts use); NO (Studey); Unclassified; DN (Gymthetic proparation); DN (Therapouts use); NO (Studey); DND (Frequested); ONS (Gymthetic proparation); ONS

ANNHER 72 OF 76 CAPLUS COPYRIGHT 2009 ACS ON STN

247067-92-1 CAPLUS 4-Thiarolidinom, 5-||5-|2,6-dichloro-4-(triflworomethyl)phenyl]-2-franylphenylese|-3-methyl-2-thioxo- (CA INDEX SUME)

93 247067-93-2 CAPLES CS 4-Thiazolidinose, 5-[(5-(4-methosphenyl)-2-Duranyl]methyleme]-3-methyl-2-thioso- (CA INDEX NAME)

24T067-95-4 CAPLUS 4-Thiarolidinome, 5-[[5-[3,5-bix(trifluoromethyl)phemyl]-2-fixamyl]methylene]-3-methyl-2-thioxo- (CA INDEX NAME)

(Contanued)

247067-87-4 CAPLUS 6-Thiarolidinome, 5-[[5-(3-chlorophenyl)-2-feranyl]methylene]-3-methyl-2-thymno: [CA THUSE NAME)

247067-88-5 CAPLUS 6-Thiarcildinome, 5-[[5-[2-chloro-5-(trifluoromethy1)pheny1]-2-furanyl]nethylens[-3-methy1-2-thioxo- (CA INDEX NAME)

247067-90-9 CAFLUS 4-Thiarolidinose, 3-methyl-2-thioxo-5-[[5-[2-(trifluoromethyl)phenyl]-2-furanyl]methylene]- (CA INDEX NAME)

PRI 247067-91-0 CAPLUS
CR 4-Thiazolidinome, 3-methyl-2-thioxo-5-[[5-[2-(trifluoromethoxy)phenyl]-2-fuxnyl]methylene]- (CA INDEX ROME)

ANSWER 72 OF 76 CAPLUS COPYRIGHT 1989 ACS on STN

247057-98-7 CAPLUS 2,4-Thiasoldidadione, 5-[[5-(4-chloropheny1)-2-furany1]methylene]-3-methyl- (CA INDEX ROME)

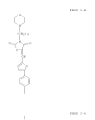
247068-00-4 CAPLUS 4-Thiasolidimone, 3-methyl-2-thioxo-5-[[5-[4-(trifluoromethyl)phenyl]-2-furanyl]methylene] (CA INDEX NAME)

247068-04-8 CAPAUS Benzenesulfonanide, 4-[5-[(3-methyl-4-oxo-2-thioxo-5-thiazolidinylideme)methyl]-2-fuzanyl]- (CA INDEX NAME)

RN 247068-06-0 CAPLUS CN 4-Thiazolidinome, 5-[[5-(4-chlorophenyl)-2-furanyl]methylene]-3-(2-

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PAGE 2-A

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HN 247068-21-9 CAFLUS CN 6-Thiarolidimone, 5-[[5-(6-chlorophenyl)-2-Curanyl]methylene]-3-[3-(18-

LT ANSWER 72 OF 76 CAPLUS COPYRIGHT 2009 ACS on STN (Continued

CB 4-Tharolidinese, 5-(5-(4-chlorophenyl)-2-furaryl)methylese)-3-(3-(4-chlorophenyl))propyl]-2-thioxo-, methanesulfonate (ht) (CA INDEX NAME) CM 1

CRS 241048-09-3 CMF C21 H21 C1 N2 C3 S2 720E 1-A

CRR 75-75-2 cleff C R4 03 S

Ris RCT (Deactant); RACT (Reactant or reagent) [preparation of thiazolidines as sisily) Lewis X synthesis inhibitor: NO 6512-99-8 CAPURE (STATE OF THE PROPERTY OF THE PROPE ARSMER TO OF TO CAPLUS COPYRIGHT 2009 ACS on STIN

Double bond geometry as shown.

226771-83-8P 226771-22-1P 226772-23-5P 226772-83-5P 226772-23-6P 226772-23-6P 226772-23-6P 226772-43-9P 226772-43-9P 226772-43-9P 226772-43-9P 226772-43-9P 226772-23-5P 22677

logical study, workszified); EMM (Synthetic preparation); TEU (Therapevtic use); ETOL (Ekological study); FAUD: (Preparation); USES (Uses) [preparation of 5-furfurylidene-4-thianoladinomes and analogs as

rular endothelial growth factor receptor amtagonists)
216771-83-4 CAPLUS
Benzensulfotanide, 4-(5-[(4-oxo-3-(phenylmethyl)-2-thloxo-5thlaroladinylidene|nethyl)-2-furanyl)- (CA INDEX NAME)

NN 216772-12-2 CAPLUS
CN Beniceses/Ifonanido,
4-[5-(]3-(],3-benicoliosul-5-ylmethyl)-4-oxo-2-thioxo-5thiarolidinylideme]methyl]-2-Euranyl]- (CA INDEX NUME)

L7 ANSMER 73 OF 76 CAPLUS ACCESSION NUMBER: 199 APLUS COPTRIGHT 2009 M/S on STN 1998:799977 CAPLUS 130:38375 Preparation of 5-furfurylideme-6-thiazolidinones and

- waveler methodising over the control of the contr

| Description | Control | State | Description | Descriptio

MARIAT 130:38375 OTHER SOURCE(S):

Title compds. [I; R = TB4; X = O, S, CR5:CR6; Rl-R6 = H, opeloalkyl, heterocyclyl, aryl, etc.; R4 = H, (opelo)alkyl, heterocyclyl, aryl, etc.; T = bond, alkylese, (alkyllimine, NBCO, etc.; Y = O, S, (alkyllimine,

CSE, E = CHE, CO, CS) were prepared as wascular endothelial growth factor receptor antagonists (no data). Thus, 3-benryl-4-thiarolidinone was acylated by Ne 5-phenyl-2-furoate (preparation each given) and the product convexted in 2 steps to I (R = CH2Ph, R1 = Ph, R2 = R3 = H, X = \circlearrowleft , Y = S,

ANSWER 73 OF 76 CAPLUS COPYRIGHT 2009 ACS on STN

216772-23-5 CAPLUS
Bemsensulfomanide, 4-[5-[[3-[[4-(dimethylamino)phenyl]methyl]-4
thioxo-5-thianolidinylidene]methyl]-2-furanyl]- (CA INDIX NUMC)

PAGE 1-A

- LT AREMER 73 OF 76 CAPLOS COPPRIGHT 2009 ACS on STN (Continued)
 CD Benzenzulfonanude,
 4-13-[12-[4-enethyl=1,3-benzodioxol-5-yl)nethyl]-4-oub-
- 232 216772-12-6 CAPLOS CB Degreesus[fosamids, 4-[5-[[3-[(3-hromopheny])methyl]-6-oxo-2-thioxo-5-thiarolidinylidens[methyl]-2-furanyl]- (CA INDEX NAME)

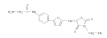
- 80-CH2-C-NH
- 300 216772-72-4 CAPLUS CP 4-Thiarolidinoss, 5-[5-(4-bromophenyl)-2-furanyl]methylene]-3-[3-(4-methyl-1-piperalmyl)-3-oxopropyl)-2-thioxo- (CA INDEX NAME)
 - PAGE 1-A

PAGE 1-A

17 AMEMIER 73 OF 76 CAPLUS CONTRIGHT 2009 ACS on STN (Continued)



381 216772-78-2 CAPLES CB Acetanide, 2-unix-8-14-[5-[14-oxo-2-[phenylmethyl]-2-thioxo-5thiazolidinylidene]methyl]-2-turanyl[phenyl]-, hydrochloride (i:1) (CJ INDEX BME)



- # 3216772-46-2 CAMARS
- 80- (CH₂)₆-NN-
- 38 216772-69-9 CAPLUS
 CN Acetanide, 2-hydroxy-N-[4-[5-[[4-oxo-3-(phenylmethyl)-2-thioxo-5-thiarolidinylideme]methyl]-2-furanyl]phenyl]- (CA INDEX NAME)
- L7 ANSWER 73 OF 76 CAPLUS COPYRIGHT 2009 ACS on STN (Continued)
 PAGE 2-A



- EN 216772-76-8 CAPLUS
 CN Benzanide,
 N-[5-[5-[6-(4-bxosombenyl)-2-furanyl]methylene]-4-oxo-2-thioxo-3-
- 288 216772-80-4 CAPLOS
 CM Acetanide, 2-anizo-8-[4-[5-[[3-[(6-methyl-1,3-benzodioxol-5-yl)methyl]-6-oxol-5-thioro-5-thioro-1dene]inethyl]-2-furanyllphenyl]- (CA INDEX
- R217-CH2-C-100
 - 221 216772-84-8 CAPLUS
 CR Renzencesfrom.nide, 4-[5-[[3-[(5-hydroxyphenyl)methyl]-4-ozo-2-thiozo-5-thiozofdisylidesejnethyl]-2-Eucasyl)- (CA 188EX 188E)

PAGE 1-A PAGE 1-A PAGE 2-A 216772-93-9 CAPLUS 4-Thiarolidinome, [4-(2-aninosetyl)-1-piperarinyl]-2-oxosethyl]-5-[[5-[4-krosophenyl)-2-furanyl)methylene)-2-thioxo- (CA INDEX NOME) PAGE 1-A PAGE 1-A

RN 216772-99-5 CAPLUS
Senzemaniforamide, 4-[5-|[3-|[3-chlorophenyl]methyl]-4-oxo-2-thioxothioxolythynichteniferbyll-2-foramil- (CA THINK MAME)

LT ARSMER 73 OF 76 CAPLUS COPYRIGHT 2009 ACS on STN

216773-47-6 CAPLUS Glycine, B-||5-||5-|4-bromophenyl)-2-faranyl]methylene]-4-oxo-2-thioxo-3-

AMSNER TO OF TO CAPLUS COPYRIGHT 2009 ACS on STN (Continued)

II 1874-89-9 18776-99-8
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**control of Control (Control of Control of Co

216774-96-8 CAPLUS 3-Thiarolidinescetic acid, 5-[[5-[4-bromophenyl]-2-furanyl]methylene]-4-am-2-thiaron- (CA NYBER NAME)

216714-19-4P 216774-28-69 Na: NCT Descript): SPM (Symthetic proparation); FRIP (Proparation); NACT (preparation of 3-furfurylidene-4-thiarologizonez and analogs as what

L7 ANSMER 73 OF 76 CAPLUS COPYRIGHT 2009 ACS on STN thiazolidinyl]acetyl]- (9C1) (CA INDEX NUME)

216775-92-3 GAING -Arginine 5-[15-(4-brosophenyl)-2-faranyl]methylene]-4-ozo-2-thioso-3-thiarolidinyl]acetyl]- (9Cl) (CA INDEX NAME)

NB 216773-61-4 CAPLOS
CR Bentemerifonantés,
4-[5-[13-[4-stro-1,3-benordioxol-5-y1]nethyl]-4-oxo2-thloxo-5-thlarolidinylidene]nethyl]-2-furanyl]- (CA IMDEX NAME)

216773-76-1 CAFLUS 4-Thiarolidinose, 5-[5-(3-hydroxyphenyl)-2-furanyl]nethylene]-3-(shenylmethyl)-2-thioxo- (CA INDEX NAME)

L7 ANSWER 73 OF 76 CAPLUS COPYRIGHT 1009 ACS on STN PAGE 1-A

PAGE 2-A

LT AREMER 73 OF 76 CARLUS COPFRIGHT 2009 ACS on STN (Continued)
REFERENCE COUNT: 3 TERES ARE 3 CITED REFERENCE AVAILABLE FOR THIS
RECORD. ALL CITATIONS WALLABLE IN THE RE

LT AREMER 74 OF 76 CAPLUS COPYRIGHT 2009 ACS on STN ACCESSION NUMBER: 1980:69311 CAPLUS DOCUMENT NUMBER: 92:69311 ORIGINAL REFERENCE NO.: 92:11209a,11292a

Data on acute toxicity of some 2-thion-3-isonicotinoylaminothiazolid-4-one

2-thion-3-isonicotinoplanisothiarolid-4-ore derivatives Danila, G., Coviereane, Bodice, Inst. Med. Farm., Lass, Rom. Bevista Medico-Chirurgicala (1579), 07(1), 121-5 CODRI, BRUIRE, 103N: 0300-0738 Southal CORPORATE SOURCE:

AB The LD50 values of 20 title tuberculostatics I (R = benrylidene, substituted benroylidene, cyclohexylidene, furfurylidene, 2-ono-2-infolene, etc.) were determined by i.p. odministration in sice.

The design of the state of the

LT ANSWER 74 OF 76 CAPLUS COPYRIGHT 2009 ACS on STN

- A8 The title thedanines I (R = Me, e-maphthyl, e-maphthylmethyl, 4-XLCGHY A1 = R, Cl, Br, Me, MeCO, MeO, ECO, ECOZO condemsed with 5-(4-nitrophemyl)- and 5-(4-nitrophemyl)thiolfurfural to give 18 corresponding I (R as above, 2 = single bond, 5) in 35.2-70.38 yield. Second-order rate constr. Cor the process and IR and OV spectral dat for

LT ARRESTS 75 OF 76 CAPLUS COPYRIGHT 2009 ACS on STN

LT ANSWER TO OF TO CAPLUS COPYRIGHT 2009 ACS on STN

PAGE 2-A

Sistems during, of the stile compode, I D = benegitions, restricted hemsyllations, or of 1.0 c. 10.0 c. 0.5 c. 0.5 c. and 1 april mere conjusted anglest Rypoductarium tobarculosis human type BITAV. The activity depends on the nature of the relieful air postulos 5 of 1.1 mg. and 1.0 c. 1.0

5-[4-(dimethylamino)bennylideme]-3-(isomicotimoylamino)-2-thiothiarolidin-4-one [1908-97-0] at 1 mg/mL were as effective as isomimid. T 6971-97-0] EL-80C (Biological activity or effector, except adverse); BSU Biological

togical study, unclassified); TBU (Therapeutic use); BIOL (Biological study);

USES

[Usas]

[Ush | (tuberculostatic activity of)

[86 69711-03-5 CAFIJS

CS 6-Fyridinecarboanide,
N-[5-[5-6-fbrosephenp]]-2-furanyl]methylene]-4-oxo2-thloxo-3-thlarolidinyl]- (CA DEDEX NUME)

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FILE 'REGISTRY' ENTERED AT 08:18:08 ON 05 MAR 2009

L1 STRUCTURE UPLOADED

L2 50 S L1

L3 7216 S L1 FULL

FILE 'CAPLUS' ENTERED AT 08:19:38 ON 05 MAR 2009 L4 192 S L3

FILE 'REGISTRY' ENTERED AT 08:28:49 ON 05 MAR 2009
L5 STRUCTURE UPLOADED
L6 4152 S L5 FULL SUB=L3

FILE 'CAPLUS' ENTERED AT 08:29:36 ON 05 MAR 2009 L7 76 S L6

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COST IN U.S. DOLLARS
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ring nodes :
1 2 3 4 5 10 11 12 13 14
ring/chain nodes :
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chain bonds :
1-6 2-7 3-8 5-15 11-15
ring bonds :
1-2 1-5 2-3 3-4 4-5 10-11 10-14 11-12 12-13 13-14
exact/norm bonds :
1-2 1-5 1-6 2-3 2-7 3-4 3-8 4-5
exact bonds :
5-15 10-11 10-14 11-12 11-15 12-13 13-14
isolated ring systems :
containing 10 :
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G1:C,O,S,N

Match level: 1:Atom 2:Atom 4:Atom 5:Atom 6:CLASS 7:CLASS 8:CLASS 10:Atom 11:Atom 12:Atom 13:Atom 14:Atom 15:CLASS

L8 STRUCTURE UPLOADED

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Structure attributes must be viewed using STN Express query preparation.

=> S L8 FULL SUB=L3

G1 C, O, S, N

FULL SUBSET SEARCH INITIATED 08:36:30 FILE 'REGISTRY'
FULL SUBSET SCREEN SEARCH COMPLETED - 7123 TO ITERATE

100.0% PROCESSED 7123 ITERATIONS 7089 ANSWERS SEARCH TIME: 00.00.01

L9 7089 SEA SUB=L3 SSS FUL L8

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This file contains CAS Registry Numbers for easy and accurate substance identification.

=> S L10 L11 13 L10

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111 ARRIMER 1 OF 13 CAPLUS ACCESSION NUMBER: 20 DOCUMENT NUMBER: 14 LUS COPYRIGHT 2009 ACS on STM 2008:1117665 CAPLUS 145:455002 163:455002 Preparation of thearolidinopse as polo like kupase inhibitors Priem, Olaf; Bohulze, Volker; Kis, Knut; Mortnarm, Lars; Kozenund, Birk; Szemzster, Gerhard; Eberspaccher, twey Gember, Judith Erittain, E. A. Schering A.-G., Germany Ger. Offen., 37pp. CODEN: GWEEK Patent Ourman PATERIT ASSIGNALI(S): DOCUMENT TYPE: LANGUAGE: FAMILY NOT. NUM: COUNT: PATENT INCOMATION: PATENT NO. | STATEST | Col. CN 101208336 PRIORITY APPIN, INFO.: US 2005-676948P P 20050503

MARRAT 145+455002

111 ANSWER 1 OF 13 CAPLUS COPYRIGHT 2009 ACS on 979

OTHER SOURCE(S):

MO 2006-894225 M 20060424

(Continued)

L11 AMEMER 1 OF 13 CAPLUS COPYRIGHT 2009 ACS on STR (Continued)

Title compdx. I $\{Y=Q(\lambda)(\lambda)\}$ Q=heteroxryly λ_s $\lambda=0$, halo, OH, etc.; = alkyl, cycloalkyl, allyl, etc.; E2 = B, halo, CB, etc.; X = NB, NE5; E5 = halo, CB, CH, etc.] and their pharmacentracilly acceptable salts were prepared For example, N-acylation of 2,2,2-triflororethamsine and carboxylic acid II [X = OB] afforded anide II [X = NBCENCTS] in 694

responsive and reliable ("Of directed under 11 in " MERALY) is an expectable published below a large of the state of the s

REFERENCE COUNT: 12 THERE ARE 12 CITED REPERENCES AVAILABLE FOR ERCORD. ALL CITATIONS AVAILABLE IN THE RE DODMAG.

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51 A2 2	20050512 90	2004-0935795	20041028
51 A3 2	20051006		
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CO, CR, CU, CE,	DE, DK, DM, DE	, EC, EE, EG,	ES, FI, GB, GD,
GH, GM, HR, HU,	ID, IL, IN, IS	JP, KE, NG,	KP, KB, KE, LC.
LR, LS, LT, LU,	LV, MA, MD, MG	, MOI, MOI, MM,	MX, ME, MA, NI,
NE, CM, PG, PH,	PL, PT, RO, RU	, SC, SD, SE,	SG, SK, SL, SY,
TN, TN, TR, TT,	TE, UN., UG, US	, DE, VC, VN,	YU, IA, IN, IN
GH, GN, KE, LS,	MW, NE, NA, SD	, SL, SI, TI,	UG, EN, EW, AN,
BY, EG, EE, MD,	BU, TJ, TN, AT	, BE, BG, CH,	CY, CE, DE, DK,
ES, FI, FR, GB,	GR, HU, IE, IT	, LU, MC, NL,	PL, PT, BO, SE,
SK, TR, BF, BJ,	CF, CG, CI, CM	, GA, GN, GQ,	GW, ML, MB, NE.
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IRFO.:		2003-514951P	

US 2003-526726P P 20031203

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compns. of the invention. Preparation of selected rhodanine compds. is

Searched by Jason M. Nolan, Ph.D.

THERE ARE 9 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE

L11 ANSMER 3 OF 13 ACCESSION NUMBER:

treatment of Pinl-associated diseases, including

DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. CO PATENT INFORMATION: PATERT NO. KIND DATE APPLICATION NO.

| PATRICUE ID. | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | 1000 | BY, BE, CA, CB, ES, FI, GB, GE, KF, KE, EE, LC, MC, MC, NA, NI, NG, SE, SE, SY, YU, IA, IN, IN IN, IN, AN, AN, CE, DE, DE, EE, FI, RO, SE, SI, BE, BW, EE, EG, KE, EG, MN, MM, SD, SE, VC, VN, TE, OG, CB, CY, NL, PL, OQ, GW,

US 2003-463271P

R DOWCE(S): MARRAT 141:376382 The invention is directed to modelators, e.g., inhibitors, of Pin1 and Pin1-related proteins and the use of such modulators for treatment of associated states, e.g., for the treatment of cancer. The present tion aims to provide photochemotherapeutic compds, with increased specificity

James and to provide photochemochemogenetic compds, with increased specificis as emposed with homes agents.

1 (7700-14-0 (7700-13-4 (7700-13-1)) (7700-14-1) (770

ANSWER S OF 13 CAPLUS COPYRIGHT 2009 ACS on STN

676651-71-1 CAPLUS 3-Thiazoldimedoutanois acid, (5-methyl-2-furanyl)setylideme)-4-oxo-2-thioxo- (CA INDEX NAME)

677000-84-9 CARUS 3-Thiazolidinebutamoio acid, 2-methyl-5-phenyl-3-Euranyl)methylene]-4-cwy-2-Ehroso- (CA IMMEX NAME)

L11 ANSWER 3 OF 13 CAPLUS COPYRIGHT 2009 ACS on STN

3-Thiazolidinebutanoic acid, -[(2-hydroxy-5-phenyl-3-furanyl)methylene]-4-oxo-2-thioxo- (CA INDEX NAME)

677002-30-1 CAPLUS

THERE ARE 4 CITED REFERENCES AVAILABLE FOR RECORD. ALL CITATIONS AVAILABLE IN THE RE

L11 AMENDER 3 OF 13 CAPLUS COPYRIGHT 2009 ACS on STN

L11 ANSMER 4 OF 13 CAPLUS ACCESSION NUMBER: 20 2004:430800 CAPLUS 140:423667 140:422667 A preparation of rhodanise derivatives, useful as inhibitors of ubiquitisation Singh, Fajander, Famesh, Usba V.; Goff, Dane, Laidig, Guy, Issalam, Sarkis D.; Hoang, Jianing; Esyan, Domald G. Domald G. Eugel Pharmscewticals, Inc., USA PCT Int. Appl., 71 pp. COURNI PICKER Patent English 4 PATENT ASSIGNACE(S): DOCUMENT TYPE: PATERT NO. APPLICATION NO US 2003-514951P P 20031028 MO 2003-0836747 W 20031113 NARPAT 140:423667 OTHER SOURCE(S):

* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT * This invention describes thodacine derive, of formula I (wheream A is 1800 to 1800 to

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NAMERA 6 Of 31 CAPAUS COMPANION 2009 AGE on GPM (Lose inseed) subshittent of biographications. The compet. and compare of the livestic are useful as inhibitions of the blookens, pathways of organizam is which biographication is in unrowlend. The invention compais, were secreted in CMP assay incarries the attachment of biographic to p30) and ANC-1/ANC-2 compared to the compared of the compared of
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|91-92-2 CAPL/S maxeladamone, 5-[[5-(2-ohlorophenyl)-2-fuxanyl]methyl]-3-(2-pxopen-1---hanne [Ch 1705K NMMS)

colating compounds and methods of use for the contains of Tail-asseminted disease, solidate manner. Books, Thomby, Debt, Soley, Hot, T., Tabletta, Thomasy, Disease, Dannase et al., Dec., USA
COMMENT PLONGS of Edge. INVENTOR(S):

PATENT ASSIGNEE(S):

DOCUMENT TYPE: LAMOUNGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

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PATERT NO.
| Description | 
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                WO 2003-086675
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OTHER SOURCE(5): MARRAT 140:315042
AB The invention is directed to modulators, e.g., inhibitors, of Pin1 and
Fun1-related proteins and the use of such modulators for treatment of

associated states, e.g., for the treatment of cancer. Synthetic methods

10/534,919 03/05/2009 L11 AMEMEN 5 OF 13 CAPLUS COPYRIGHT 2009 ACS on STR

(Continued)

LLI ARSMER 5 OF 13 CAPLUS COPYRIGHT 2009 ACS on STR

NN 676651-71-1 CAPLON
CN 3-Thiarolidinebutanord acid,
5-[3-(5-methyl-2-furanyl)butylidene]-4-ozo-2thioco- (CA INDEX NAME)

322 677000-04-2 CAPLUS
CH 2-Thisplathebutanoid acid,
5-[(2-methyl-5-phenyl-3-furanyl)methylene]-4oxo-2-thioxo- [CA IMDEX NAME)

FORMAT

PR 677001-10-4 CAPLOS CR 3-Thiasolidisebutanosc acid, 5-[(2-hydroxy-2-phesyl-3-furanyl)methylene]-4-oxo-2-thioxo- (CA INICK NAME)

677002-30-1 CAPLOS 3-Thiarolidimebutamoic acid, 5-(3-furamylmethylene)-4-omo-2-thiomo- (CA NEDEX NUME)

LL1 ANSWER 5 OF 13 CAPLUS COPYRIGHT 2009 ACS on STN (Continued) (CH2)3-CO28

LL1 ANSMER 6 OF 13 CAPLUS COPYRIGHT 2009 ACS on STN
ACCESSION NUMBER: 2000/184390 CAPLUS
TITLE: Preparation of arylmethylene
heteroryclylmethylene

thiarolidimediones and analogs as tumor necrosis factor inhibitors Wang, Jing; Rammarayan, Kalyanaraman; Rideout, Mong, Seymour; Zhu, Bengy; Niemeyer, Christins; Brady, Thomas P. Stroctural Bioinformatics Inc., USA NCT Int. Appl., 127 pp. CODER: FIXE2

PATENT ASSIGNEE(S): Patent English DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

PATERT NO. MIND DATE APPLICATION NO.
A1 20000608 MO 1999-0528856 US 1999-316415 A 19990521

OTHER SOURCE(S): MARPAT 133:30722

AB The title compds: (I) [wherein WI-W5 together = aliphatic, heterocyclic,

or betroarco. Indep 31 = N or (mo)substituted heterosyclia, (betroalcontically) 21 = N or (N)substituted heterosyclia, (betroalcontically) 21 = N or S) and sualogs were prepared by condensing almosystem with this holizable decorate. For example, the substituted is a substituted by the substitute by the subst

data). 1100394-27-1 EL: FFSH (Prophetic) (Preparation of arylmethylene and heterocyclylmethylene

L11 AREMER C OF 13 CAPLIS COPPRIGHT 2009 ACS on STN (Continued) thisiolidinediones and analogs as twoor necrosis factor inhibitors) 39 1200394-27-1 CAPLIS C PROPERTY C PROPERTY

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L11 ANSMER 7 OF 13 CAPL ACCESSION NUMBER:

nate and the appendix drugs containing the continues that the appendix drugs containing the continues that the poly: Barakane, Baltimor Tatest, Allias Bankalis, Marias Bankalis

PATENT ASSIGNED (S):

DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

PATERT NO. KIND DATE APPLICATION NO. DATE JP 2000095770 PRIORITY APPLES, IMPO.,

OTHER SOURCE(S): NARPAT 132:246365

AB The title derivs. I [Y = {un}substituted aryl, {un}substituted nono or condensed heterocyclylr X = sulfonyl, carbonyl, carbonylony, thiocarbonylony, W = O, Sp. Z = {un}substituted aryl, {un}substituted

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2. ANNERS 1 OF 13 CAPTED CONTRIGUET DOD ACE on STM (Constrained) SIGOL (Babological study), PSEP (Preparation), DIESE (Sees) (obypase inhibitors for treatment of cardiovascular diseases osused by 12 (2012) (Constrained Constrained Constrained

energy="10"0" --enrol5 2,4=Thiazolidimediome, 5-(3-furamylmethyleme)-3-(5-quimolimylsulfomyl)-(CA INDEX NAME)

929 262602-76-6 CAPLUS

ANSWER 7 OF 13 CAPLUS COPYRIGHT 2009 ACS on STM (Continued) 2.4-Thiszolidinedione, 3-(3,4-dimethoxybenzoyl)-5-(3-Euranylmethylene) (CA INDEX NAME)

CN 2,4-Thiarolidizedione, 5-(3-furanylmethylene)-3-(2-naphthalenyloarbomyl)-

262802-78-8 CAPLUS 3-Thiarolidimecarboxylic acid, 5-(3-furanylmethylene)-2,4-dioxo-2-maphthalonyl ester (CA INDEC NAME)

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Lil AMEMBER 8 OF 13 CAPLUS COPYRIGHT 2009 MCS on STM (Continued)
AB Compdr., comput. and methods are provided for the treatment and
prophylaxis of infections and associated diseases caused by viruses of
   L11 AMENDER 8 OF 13 CAPLIES ACCESSION NUMBER: 20
                                                                                                                                                                                                                                                                                                                                                                                       prophylatic of infections and associated diseases council by virtues of the Turbivitizing Canally, by edistricting because products on the control of the Canal Ca
                     PATERT NO.
RN 1100514-41-7 CAPLUS
CN 3-Thianolidineacetic acid,
5-[[5-(4-carboxypheny1)-2-furany1]methy1]-4-oxo-
2-thioxo- (CA INDEX NOME)
                                                                                                                                                                              US 1998-113212P
                                                                                                                                                                                                                                                           P 19981222
                                                                                                                                                                              US 1999-119328P
                                                                                                                                                                                                                                                                                                                                                                                         3-Thiazolidinepropanoic acid, 4-oso-5-[[5-(2-phenylethynyl)-2-
furanyl methyl-2-thioso- (CA INDEX NUME)
                                                                                                MARPAY 132:189652
   111 ANSWER 0 OF 13 CAPLUS COPYRIGHT 2009 ACS on S78
                                                                                                                                                                                                                                                                                                                                                                     LII ANSWER 0 OF 13 CAPLUS COPYRIGHT 2009 ACS on STN
                                                                                                                                                                                                                                                                                                                                                                                       1100515-07-8 CAPLUS
3-Thiarolidinebutamoic acid,
5-(3,4-dichlorophemy1)-2-furamy1]methy1]-
4-xxx-2-thixxx- (CA INDEX NUME)
     PR 1100514-47-3 CAPLUS

CN 3-Thiarolidinepropanels acid,

4-cxc-5-[[5-(2-thiaroly1)-2-furany1]methy1]-

2-thicxc- (CA INDEX NAME)
                                                                                                                                                                                                                                                                                                                                                                                         3-Thiazolidinepropanoic ecid, 4-oxo-5-[(5-pheny1-2-furany1)methy1]-2-thioxo- (CA INDEX NUME)
                       1100515-05-6 CAPLUS
3-Thiarolidinepropanoic acid, 5-[[5-(3-chloropheny1)-2-furany1]methy1]-4-
our-2-thorop- (CA TMIDE NAME)
                                                                                                                                                                                                                                                                                                                                                                                         CH2-CH2-CO2H
                                                                                                                                                                                                                                                                                                                                                                                     1100515-12-6 CAPUS
3-Thiazolidineacetic acid,
co-5-[(5-phenyl-2-furanyl)methyl]-2-thico-
(CA INDEX NAME)
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LLI ARSMER 8 OF 13 CAPLUS COPYRIGHT 2009 ACS on STR

THERE ARE 6 CITED REFERENCES AVAILABLE FOR THIS RECORD, ALL CITATIONS AVAILABLE IN THE RE TORRAT

L11 ANSMER 9 OF 13 CAPLUS ACCESSION NUMBER: 19

LUS COPYRIGHT 2009 ACS on STN 1999:699310 CAPLUS 131:299442 131,292442
Preparation of this rollidings as sualyl Lewig X
Preparation this bittors
Poblayashi, Koruy Nishuyana, Toshikhikoy Nakaide,
Shanja,
Shanja, Cook Cook, Japan
Sps. Koka: Tokkyo Koho, 30 pp.
CODRI JOSCAN

PATENT ASSIGNMENTS:

DOCUMENT TYPE:

PATERT NO. KIND DATE APPLICATION NO DATE JP 11302280 PRIORITY APPLES, INFO.:

OTHER SOURCE(S): NARIWT 131:299442

AB The title compdx. I [Al, A2, A3 - O, S; E1 - alkyl, alkenyl, etc.; E2 alkyl, etc.; n=1-3; ring Bl=heterocyclic ring, etc.; dotted line indicates single or double bond) are prepared. In an in vitro test using <math>Bl=0 cells, the title compound II at 3 μM yave 1004 inhibition of

Statyl Lewis X synthesis. Formulations containing I are given.
17 247058-14-07 Ri RWC (Biological scrivity or effector, except adverse); BSU (Biological)

[Biological unclassified) STR (Synthetic preparation); 7EW (Therapeutic use); Biological stady); PEW (Preparation); OSES (Deep ((preparation of thistolidines as sisky) Lewis X synthesis inhibitors) 28 24765-14-0 CARUS

AMENER 9 OF 13 CAPLIS COPYRIGHT 2009 ACS on STM (Continued) 4-Thiarolidinome, 3-methyl-5-[15-[4-mitrophenyl)-2-furnyl]methyl)-2-thicore (CA INDEX NAME)

LII ANSMER 10 OF 13 CAPLUS COPPRIGHT 2009 ACS on STN
ACCESSION NOMBER: 1995:22:007 CAPLUS
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TOCKNERT WRITE 1995:22:007 CAPLUS
TITLE: 1995:22:0

ORATE SOURCE:

chodanines. Bart IV Donate, Jr. Science, Benha Gaiveraity, Benha, Egypt Egyptian Journal of Pharmacoettani Sciences (1974), Wolmen Bate 1997, 34(4-6), 521-8 COUDER EXTRESS; 1984: 0901-0904 Journal Lindonstein and Documentation Centre Doctania

Substituted rhodanines I (R = thienyl, furyl, pyrrolyl; R1 = B) reacted with halo compds. aromatic aldehydes, betomes, anhydrides, and animes to

I (add P) F1 = No, Pb, CEICEFD, etc.). The antibaterial activates of 10097-07-09ericed deriva. have been investigated. Its SECTION OF THE PROPERTY OF THE PROP

Lil AREMER 11 OF 13 CAPLUS COPYRIGHT 2009 ACS on STN ACCESSION NUMBER: 1994:134356 CAPLUS DOUMERT NUMBER: 129:134356

DOCUMENT NUMBER: ORIGINAL REFERENCE NO.:

DOCUMENT TYPE: LANGUAGE: OTHER DOUBCE(D):

Assessing of Yearly-In-Thomas-Albientinous with Neumain Colombia. Secured on For Johnson St. Colombia was when discovered in 18 or 10 to 1

111 ANSWER 12 OF 11 CAPLUS COPYRIGHT 2009 ACS on STN

5-Thiarelidineacetonitrile, a-2-furanyl-3-[3-methyl-2-[](4-methylmhemyl)sulforyllamine)-1-encentyll-4-core-2-thicore (9CI) (CA INDEX

132177-44-7 CAPLUS Benzenceulfonanido, N-[1-[(5-(oyano-2-furanylmethyl)-4-oxo-2-thloxo-3-thlazoladinyl]oarbonyl]-3-methylbutyl]-4-methyl- (CA INDEX NAME)

L11 ANSMER 12 OF 13 CAPLUS COPYRIGHT 2009 ACS on STN ACCESSION NUMBER: 1991:101806 CAPLUS DOCUMENT NUMBER: 114:101806

114:10:1006
114:17:55a,17:55a
The synthesis and biological activity of 3,5-disabitited inbolanizes. Fart II benus, Shafise G. Kec. Sci., Zapaig Guir, Benhs, Egypt Journal of the Sethian Chemical Society (1989),

407-15 CODER: JSCSER; ISSN: 0352-5139

DOCUMENT TYPE: LANGUAGE: OTHER SOURCE(S): Journal English CASEEACT 114:101806

A new class of rhodanine derivs. with phthalyl I (R = Ne, GB, CN; R1 = 2-pyryl, 2-thipphenyl, 2-furyl; E2 = Gly, Ala, Ser) and tosyl II (E2 = Gly, Val, Lee) amino acid modelies was memored. All the synthesized

derive.

When correspond for antinirobbil activity.

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12217-46-77 32246-55-99 122146-62-47

12217-46-77 32246-55-99 122146-62-47

[Biological activity or effector, except adverse); RSU (Biological activity or effector, except adverse); RSU (Biological activity or effector).

| Riological study, medissified); ERR (Symbolic preparation); EIG. (Riological study); PERP (Preparation) and individual study); PERP (Preparation) and individual study); PERP (Preparation); PERP (Preparati

ASSESSE 11 OF 13 CALING CONTRIGHT 2009 ACS on STH (Continued)
123246-79-29 TRANSPORT OF TREPARATION)
props. 621
props. 622
18-1281602-1,3(28)-46068, 2-[2-[5-(2-frazyllystosymethyl)-4-ozo-2-thioso-2-thiosolidyspl-2-ozo-2017)

132146-43-1 CAPLUS 13-1soindole-1, 3(22)-dione, 2-{2-{5-{42-furanylhydroxymethyl}-4-oxo-2-thioxo-3-thiarolidlinyl}-1-thydroxymethyl}-2-oxoethyl}- (CA INDEX NNE)

132146-44-2 CAPLUS Benzenezulfonanide, N-[2-[5-(2-furanylhydroxymethyl)-4-oxo-2-thioxo-3-thiazolaldisvll-2-oxosethyll-4-methyl- (CA INDEX NOME)

132146-45-3 CAPAUS 4-Thiarolidimone, 5-(2-furanylhydroxymethyl)-3-[3-methyl-2-][(4-methylohenyl)zulfonyljamino]-1-oxobutyl]-2-thioxo- (9CI) (CA INDEX NAKE)

L11 ANSMER 12 OF 13 CAPLUS COPYRIGHT 2009 ACS on STN (Continued)

132146-50-0 CAPAUS 5-Thiaralidinacetonitrile, 3-[2-[1,3-dibydro-1,3-dioxo-2B-isoindol-2 yllacetyl]-a-2-furanyl-4-oxo-2-thioxo- (CA INDEX NAME)

N 132146-60-2 CAPLUS N 5-Thiarolidimaretomitrile, -[2-(1,3-dihydro-1,3-dioxo-2E-imoindol-2-yl)-3-hydroxy-1-compropy]-a-2-furanyl-4-oxo-2-thioxo- (CA INDEX NAME)

132146-61-3 CAPLUS Benzensulfonamide, N-[2-[5-(oyano-2-furanylmethyl)-4-oxo-2-thioxo-3-thiaroidinyi]-2-oxoethyl]-4-methyl- (CA_INDEX_NAME)

111 ANNUAR 12 OF 13 CAPLUS COPYRIGHT 2009 ACS on STS (Continued)

132146-77-1 CAPLUS
Benrecesulfocamide, N-[2-[5-[1-(2-furanyl)ethyl]-4-oxo-2-thioxo-3-thiarolidinyl]-2-oxoethyl]-4-methyl- (CA_INDEX_NAME)

132146-70-2 CAPLUS 4-Thiarolidinose, 5-[1-[2-furanyl)ethyl]-3-[3-methyl-2-]|[4-methylphenyl)sulfonyl]amino]-1-oxobutyl]-2-thioxo- (9CI) (CA INDEX NAME)

$$\label{eq:continuous} \begin{split} &132146-79-3 \quad \text{CAPLIUS} \\ & \text{Entraceaulforanide}, \quad N-\{1-||5-|1-(2-\text{furanyl})\,\text{ethyl}\}-4-\text{exc-}2-\text{thicxo-}3-\text{thiazolidinyl}\,\text{leatboyl}\}-3-\text{esthyl}\,\text{levtyl}\}-4-\text{exthyl}-\quad (CL INDEX NME) \end{split}$$

781 132146-75-9 CAPLOS
CB 18-Incindole-1,3(28)-dione,
2-[2-[5-[1-(2-furanyl)=thyl]=4-cuo-2-thioxo-3-thiarolidinyl]=1-methyl=2-cuoethyl]= (CA INDEX NUME)

IR 132.44- (4-0 CAPLOS
CR 18-1-301nfols-1,3(2E)-diome,
2-[2-[5-[1-(2-fmanyl)ethyl]-4-oxo-2-thioxo-3thiagolidisyl]-1-(hydroxymethyl)-2-oxo-thyl)- (CA INDEX NOME)

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are agitated 5 hrs., a solution of 0.1 mole CIJCCOLK in 20 ml. HJO added, agitation continued 1 hr., and the mixture warmed to 90° on a water bath, made strongly acidic with concentrated HCl, and cooled to give

yellow 3-furfurylrhodanime (I), m. 73-4" (MeOB), as compared with 11% by the method (Erown, et al., CM. 50, 12963a) using NEGOS rather than NOBL. A mixture of 0.0075 mole 1, 0.0075 mole appropriate aldebyde, 20

NOS. A mixture of 0.027 mais 3, 0.007 mais appropriate alonyjes, 25 Mod. and 14 g, 500 mt efficient of 3 Max. sylinded the Dissived per 1, 100 mt of 1, 100 mt of

4-Thiazolidinone, 3-(2-furanylmethyl)-5-(3-furanylmethylene)-2-thioxo-(CA INDEX NAME)